

The shrunken genome of *Arabidopsis thaliana*

Ryan K. Oyama · Maria J. Clauss · Nataša Formanová · Jürgen Kroymann ·
Karl J. Schmid · Heiko Vogel · Kerstin Weniger · Aaron J. Windsor ·
Thomas Mitchell-Olds

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Abstract This paper examines macro and micro-level patterns of genome size evolution in the Brassicaceae. A phylogeny of 25 relatives of *Arabidopsis thaliana* was reconstructed using four molecular markers under both parsimony and Bayesian methods. Reconstruction of genome size (*C* value) evolution as a discrete character and as a continuous character was also performed. In addition, size dynamics in small chromosomal regions were assessed by comparing genomic clones generated for *Arabidopsis lyrata* and for *Boechera stricta* to the fully sequenced genome of *A. thaliana*. The results reveal a sevenfold variation in genome size among the taxa investigated and that the small genome size of *A. thaliana* is derived. Our results also indicate that the genome is free to increase or decrease in size across these evolutionary lineages without a directional bias. These changes are accomplished by insertions and deletions at both large and small-scales

occurring mostly in intergenic regions, with repetitive sequences and transposable elements implicated in genome size increases. The focus upon taxa relatively closely related to the model organism *A. thaliana*, and the combination of complementary approaches, allows for unique insights into the processes driving genome size changes.

Keywords Genome size · *C* value · Brassicaceae · *Arabidopsis* · *Boechera stricta* · Phylogeny · Repetitive sequence · Transposons

Introduction

The fundamental riddle in genome size studies lies in the observations that across all of life there is large variation in genome size but low variation in the number of genes, and that genome size and phenotypic complexity seem to have little relationship with one another (Gregory 2005a; Nagl et al. 1983). This disconnect between genome size/complexity and phenotype was initially dubbed the “*C* value paradox” (Thomas 1971). Efforts to explain this conundrum have greatly increased our understanding of the genome but questions remain regarding the dominant mechanism for genome size change and the evolutionary forces that may be operating (Bennett and Leitch 2005b).

It has long been clear that dramatic increases in plant genome size can be accomplished via polyploidization (Bennetzen 2000; Bennetzen et al. 2005; Vitte and Panaud 2005; Wendel 2000). However, the lack of obvious mechanisms by which reductions in genome size could be accomplished led to a suggestion that plants have a “one-way ticket to genomic obesity” (Bennetzen and Kellogg 1997). Since then, potential mechanisms by which genome size can increase or decrease, especially via small and

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R. K. Oyama (✉) · M. J. Clauss · N. Formanová · J. Kroymann ·
K. J. Schmid · H. Vogel · K. Weniger · A. J. Windsor ·
T. Mitchell-Olds
Max Planck Institute for Chemical Ecology,
Hans-Knoell Strasse 8, 07743 Jena, Germany
e-mail: royama@lrz.uni-muenchen.de

Present Address:

R. K. Oyama
Systematic Botany, Ludwig-Maximilians University,
Menzingen Str. 67, 80638 Munich, Germany

Present Address:

T. Mitchell-Olds
Department of Biology, Duke University, PO Box 91000,
Durham, NC 27708, USA

gradual changes, have been identified and demonstrated (Devos et al. 2002; Petrov 1997). Nevertheless, it remains unclear if these mechanisms are actually responsible for the changes observed across evolutionary time.

The small size of the *A. thaliana* genome ($n = 5$, $C = 0.16$ pg), among the smallest published (Bennett and Leitch 2005a), has made it a favored system in which to investigate questions of gene regulation and function, and genome organization. This has revealed a complicated history of genome and chromosome level duplication and rearrangement in *A. thaliana* as illustrated by Lysak et al. (2006) (for a review see Schranz et al. 2007). It is clear that simply adding together the micro-level processes will not necessarily provide us with a full understanding of genome evolution (Charlesworth et al. 2001; Gregory 2004, 2005b). If we are to have a realistic understanding of general processes in genome evolution then we must branch out from “model” species (Gregory 2005a; Koch and Kiefer 2005). Thus, greater clarity might be leveraged by analyzing the genomic data we now have for *A. thaliana* within a comparative framework that includes its relatives.

The family Brassicaceae has several attributes that make it attractive for the study of genome size evolution. The family is large, with ca. 340 genera and 3,700 species (Al-Shehbaz et al. 2006), has great variability in life history, habit, and ecology, and is known to have undergone recurrent polyploidization over the course of its evolution. The family displays a moderate diversity of genome sizes, (Johnston et al. 2005; Nagl et al. 1983) and a range of monoploid chromosome numbers (Warwick and Al-Shehbaz 2006). Finally, there is a considerable amount of genome level information already available from multiple genera and species within Brassicaceae in addition to *A. thaliana*. These include emerging model species such as *Arabidopsis lyrata*, *Boechera stricta*, *Brassica oleracea*, and *Capsella rubella*.

Previous studies have begun to untangle the phylogenetic relationships within the Brassicaceae (Beilstein et al. 2006; Hall et al. 2002, 2004; Koch et al. 2001; O’Kane and Al-Shehbaz 2003; Warwick et al. 2002). A few studies have also approached the evolution of genome size within a phylogenetic context (Bennetzen and Kellogg 1997; Jakob et al. 2004; Soltis et al. 2003; Weiss-Schneeweiss et al. 2006). However, only one study has focused on genome size evolution among the taxa near to *Arabidopsis thaliana* (Johnston et al. 2005). But, it relied on only one marker for the reconstruction of the phylogeny and did not apply significance tests to the evolution of the character. Furthermore, no study has attempted to investigate the micro-level patterns that might reveal the actual mechanisms responsible for genome size changes among these evolutionary lineages. Thus, there is an opening for a study that

combines a solid phylogeny of species related to *A. thaliana* with reconstruction and testing of genome size evolution upon that phylogeny and an investigation of possible mechanisms.

Here, we present the results from a comparative investigation into the evolutionary history of the small genome of *A. thaliana* that is centered around a robust and well-sampled phylogeny recovered using four genes and sampling of the genera thought closely related to *Arabidopsis* (Al-Shehbaz et al. 2006; Beilstein et al. 2006; Koch and Kiefer 2005). The character mapping of genome sizes onto this phylogeny thus gives a broad level view into the pattern of genome size evolution among *A. thaliana* and its close relatives. In addition, we delve into the patterns and mechanisms underlying the change in genome size. This is accomplished by comparing end-sequences of genomic shotgun clones from *A. lyrata* (O’Kane and Al-Shehbaz 1997) and *B. stricta* (Al-Shehbaz 2003) against the fully sequenced genome of *A. thaliana*. This combination of approaches allows for a robust analysis of genome size evolution in this area of Brassicaceae.

Materials and methods

Taxon sampling

We included 26 terminal taxa representing 23 genera of Brassicaceae (see Table 1), including as many genera as possible from the Cardaminoid, and Halimolobine genera near *Arabidopsis* (Koch et al. 2005). We used sequences from *Aethionema* as an outgroup based on the availability of sequences and the use of this taxon as an outgroup by other studies of this area of Brassicaceae (Bailey et al. 2002; Koch et al. 2001).

DNA isolation, amplification, and sequencing

Total genomic DNA was extracted from silica-gel dried material, herbarium specimens, or fresh leaf material using a modified 2× cetyltrimethylammoniumbromide (CTAB) method (Doyle and Doyle 1987) or, alternatively, using QIAGEN Genome DNA prep (Qiagen GmbH, Düsseldorf, Germany). The DNA was then used to amplify four markers by PCR under standard conditions: Chalcone Synthase (*Chs*), ITS, *matK*, and *trnL-F*. The *Chs* primers span the last exon and were the same as used by Koch et al. (1999). The primers used to amplify *matK*, *trnK-710F* and *trnK-2R*, originally derived from Johnson and Soltis (1995), were the same as used by Koch et al. (2001). The ITS primers amplified a region that included the 5.8S region and the flanking portions of the 18S and 26S regions (Baum et al. 1998; White et al. 1990). The *trnL* primers

Table 1 List of taxa included in this study along with corresponding GenBank accession numbers for each marker sequenced, the genome size (2C), the number of biological replicates used to measure and calculate genome size, and the chromosome count (2n)

Taxon	ITS	<i>Chs</i>	<i>matK</i>	<i>trnL</i>	Genome size (pg)	SE \pm	Rep.	2n
<i>Aethionema</i> ^a	AY254539	AF112082	AF144354	AY122451	0.342	–	–	22
<i>Arabidopsis halleri</i>	AJ232894	AF112095	AF144341	DQ406772	0.252	0.003	8	16
<i>Arabidopsis lyrata</i> ^b	AF137539	AF112104	AF144336	DQ406773	0.251	0.002	23	16
<i>Arabidopsis thaliana</i>	X52322	AF112086	AF144370	NC000932	0.176	0.002	3	10
<i>Barbarea vulgaris</i>	X98632	AF112108	AF144330	AF198119	0.275	–	1	16
<i>Boechera stricta</i>	DQ399111	AF112088	AF144343	DQ406776	0.262	0.002	2	14
<i>Camelina microcarpa</i>	DQ399112	DQ399095	DQ406760	DQ406777	0.277a	–	1	26
<i>Capsella rubella</i>	AJ232913	AF112106	AF144334	DQ406778	0.245	0.017	2	16
<i>Cardamine amara</i>	DQ399113	DQ399096	AF144337	DQ406779	0.225b	b	b	16
<i>Catolobus pendula</i>	AF137572	DQ399093	DQ406758	DQ406774	0.365	0.004	3	16
<i>Crucihimalaya wallichii</i>	DQ399114	DQ399097	AF144367	DQ406780	0.327	0.014	2	16
<i>Cusickiella douglasii</i>	AF146515	DQ399098	DQ406761	DQ406781	0.476	0.004	2	14
<i>Erysimum perovskianum</i>	DQ399116	DQ399099	DQ406762	DQ406782	0.352	–	1	36
<i>Eutrema halophilum</i>	AJ232925	DQ399108	AF144333	DQ406793	0.336	–	1	14
<i>Fourraea alpina</i>	AJ232890	AF112102	AF144335	DQ406783	0.478	0.008	4	14
<i>Halimolobos jaegeri</i>	DQ399117	DQ399100	DQ406763	DQ406784	0.221	0.003	3	16
<i>Lepidium perfoliatum</i>	DQ399120	DQ399102	DQ406766	DQ406787	0.304	–	1	16
<i>Neslia paniculata</i>	AF137576	DQ399103	DQ406767	DQ406788	0.213	–	1	14
<i>Olimarabidopsis pumila</i>	AY254546	DQ399104	AF144345	DQ406789	0.347	–	1	32
<i>Phoenicaulis cheiranthoides</i>	DQ399121	DQ399105	DQ406768	DQ406790	0.391	–	1	14
<i>Polycetenium fremontii</i>	DQ436937	DQ399106	DQ406769	DQ406791	0.588	0.004	8	14
<i>Rorripa islandica</i>	DQ399122	DQ399107	DQ406770	DQ406792	0.263	0.001	2	16
<i>Sandbergia perplexus</i>	DQ399118	AF112094	DQ406764	DQ406785	0.573	0.009	3	14
<i>Sandbergia whitedii</i>	DQ399119	DQ399101	DQ406765	DQ406786	0.780	0.015	2	14
<i>Sisymbrium irio</i>	AF531568	AF144541	AF144366	AY236216	0.349	0.007	2	14
<i>Transberingia bursifolia</i>	DQ399110	DQ399094	DQ406759	DQ406775	0.232	0.003	2	16
<i>Turritis glabra</i>	AJ232925	DQ399109	AF144333	DQ406794	0.227	0.006	3	12

Unless otherwise noted, genome size values are derived from measurements made for this study

^a Sequences are of *Aethionema grandiflora* except for *trnL*, (*A. saxatile*), and ITS (*A. arabicum*). Genome size is based on measurements of *A. stylosum* (E. Schranz, personal communication)

^b *Arabidopsis lyrata* subsp. *petraea*

a = Genome size is of *Camelina laxa*; b = Genome size as reported in Johnston et al. (2005)

used corresponded to *trnC* and *trnD* from Taberlet et al. (1991). All of the primers used for amplification are summarized in Table S1 in Electronic supplementary material.

All of the PCR products were directly sequenced using the same primers as for PCR with BigDye chemistry (ABI Perkin-Elmer, Foster City, California, USA). Reactions that could not be directly sequenced were first cloned and 6–12 clones were then sequenced. Under this procedure PCR products were gel-cleaned and then cloned using the TOPO TATM kit (Invitrogen Corp., Carlsbad, California, USA). Plasmid DNA was purified using Qiaquick Mini-Prep kits (Qiagen Corp.,) and then sequenced using the aforementioned amplification primers and the following internal primers for *matK*: *matK*-1010R, *matK*-1089R, and

matK-1495R. These primer sequences are also included in Table S1.

Phylogenetic analyses

Sequences were assembled and edited using SeqManTM (DNASTAR Inc., Madison, Wisconsin, USA), and manually aligned using MacCladeTM (Sinauer Associates Inc., Sunderland, Massachusetts, USA). There was little ambiguity in alignment. However, when there were alternatives that seemed equally plausible, we chose the alignment that minimized the number of potentially informative characters to avoid biasing the results. Sequences generated for this study were submitted to GenBank with accession

numbers as given in Table 1. Voucher specimens were deposited at M and DUKE.

Parsimony analyses were performed using PAUP* version 4.0b10 (Swofford 2002). All searches for most parsimonious trees were conducted with tree bisection-reconnection (TBR) heuristic searches and 100 random taxon addition replicates. Bootstrap analyses were done with 100 bootstrap replicates (Felsenstein 1985), as implemented in PAUP* version 4.0b10, with the search conditions the same as described previously. In order to test congruence between our nuclear and chloroplast markers, partition homogeneity tests were conducted in PAUP* 4.0. We used the heuristic search strategy previously described with 999 random replicates to generate the distribution resulting in the *P* values.

Bayesian phylogenetic analysis on the combined dataset was performed using Metropolis-coupled Markov-chain Monte Carlo (MC³) analysis, as implemented in the program MrBayes 3.1 (Huelsenbeck and Ronquist 2001). Given that Bayesian analysis requires a model of molecular evolution to be specified, we used the program ModelTest 2.2 (Posada and Crandall 1998) to select an appropriate model of DNA substitution. For this dataset, ModelTest 2.2 selected a general time-reversible model (GTR) with a discrete approximation to the gamma distribution for rate variation among sites (Γ) and a proportion of invariant sites (I) (Yang 1994a, b).

Tree searching using MrBayes was performed by running five coupled chains initiated from random trees (sequential heat = 0.2) for 1,000,000 generations with trees sampled every 100th generation (Huelsenbeck and Ronquist 2001). At the end of the run, convergence was evaluated by visual inspection of a graph of likelihood as a function of generation. A conservative burn-in period was identified, and only post-burn-in trees were saved. This analysis was repeated four times, and the majority-rule consensus trees from each run were compared to evaluate mixing. The four sets of post-burn-in trees were then pooled to form a majority rule consensus tree, and this pool was taken as the best representation of the posterior distribution of tree topology and model parameters (Huelsenbeck et al. 2002; Miller et al. 2002).

The phylogenetic placements of taxa that were uncertain were further examined using the Templeton test (Templeton 1983) and the Shimodaira-Hasegawa test (Kishino and Hasegawa 1989; Shimodaira and Hasegawa 1999) as implemented in PAUP* 4.0b10 (Swofford 2002) on the combined dataset. In each case, the most optimal tree was compared to a tree recovered in searches that constrained the taxon of interest to its alternative placement. The Shimodaira-Hasegawa tests assumed the GTR + Γ model parameters (Yang 1994a, b). If the *P* values were less than 0.05, the two trees were considered significantly different.

Reconstruction of genome size evolution

Genome size measurements for each taxon were obtained from fresh material of diploid individuals (i.e. we excluded polyploid individuals) using a PartecTM CCA-II flow-cytometer and CyStain UV precise P nuclei extract and staining kit (Partec GmbH, Münster, Germany). All measurements used *B. oleracea*, *B. rapa*, *Matthiola incana*, or some mixture of those three species as standards and were recorded as a ratio of the size of the plant being measured to *B. oleracea*. These ratios were converted into 2*C* values using a published value of 0.710 pg for *B. oleracea* (Johnston et al. 2005). One *C* value (*Cardamine amara*) was obtained from Johnston et al. (2005). The values we obtained agreed with those available in the *C* values Database (<http://www.kew.org/uk/cval/homepage.html>). These values and the phylogeny were used to examine the phylogenetic constraints on genome size evolution as well as its directionality, mode, and tempo.

The genome sizes (2*C*) were converted into a discrete character by grouping them into nine size ranges of 0.1-pg intervals and each interval designated as a character state. This character of genome size was mapped onto the phylogeny using MacCladeTM under ACCTRAN parsimony optimization under both an ordered and unordered treatment of the character. In order to test for phylogenetic constraint of genome size evolution, a permutation test was conducted in MacCladeTM that resembled a test of phylogenetic autocorrelation for discrete characters. To conduct this test, the character of genome size was replicated 999 times (for a total of 1,000 characters) and then the states were shuffled among the taxa. This resulted in a distribution of steps across the 1,000 characters that approximated the normal distribution with a slight skew toward longer trees. If the number of steps necessary to explain the original character was within the lower or upper 5% of the distribution, then the distribution of genome-size on the phylogeny was considered to be significantly non-random.

Analysis of genome size (2*C*) as a continuous character was done using Continuous (Pagel 1997, 1999). This program is able to test whether there is directionality to genome size changes by comparing likelihood ratio statistics recovered from a random walk model and a directional random walk model. The program also allows for the estimation of parameters (Lambda, Kappa, and Delta), which, respectively, capture the degree to which the evolution of the character is explained by the phylogeny, whether the changes occur gradually, and whether the pattern of changes is consistent with an adaptive scenario. In order to conduct a more conservative test, the values for these parameters were first estimated under Maximum Likelihood and the random walk model and then used to

conduct a Maximum Likelihood ratio test between the directional and non-directional random walk models.

The evolution of chromosome numbers ($2n$ values) across the phylogeny was also analyzed using MacClade™ and Continuous. The values were obtained from published sources (Al-Shehbaz et al. 2006; Bailey 2006; Lysak et al. 2003; Warwick and Al-Shehbaz 2006). For the MacClade™ analyses, the chromosome counts ($2n$) for each taxon were converted into a discrete character state (a range of 14 possible states) and treated as unordered or ordered as described earlier. We also performed the modified permutation test as described for genome size measurements on the character of chromosome number. The Continuous analyses used the same methods as used for genome size. We also used Continuous to test whether or not the evolution of genome size ($2C$, pg) and chromosome number ($2n$) are correlated on the phylogeny. In this case, the values of Kappa, Delta and Lamda were first optimized using both characters. Then all of the possible combinations of directional and non-directional models were compared while forcing the covariance of the two characters in the null model to be zero (i.e. complete independence).

Genomic library construction

Size-selected genomic shotgun libraries for both *B. stricta* and *A. lyrata* were constructed in the pZERO-II plasmid vector (Invitrogen, Karlsruhe, Germany). Approximately 5 µg of plant genomic DNA were partially digested with *Sau3AI* (New England Biolabs, Frankfurt am Main, Germany). The DNA was then dephosphorylated by incubation with 0.1 U of CIAP (Amersham, Freiburg, Germany), extracted once with phenol/chloroform, ethanol-precipitated, dissolved in TE, and size-fractionated by means of gel electrophoresis on a 0.7% agarose gel. DNA fragments in the size-interval from 4.5 to 5.5 kb were excised in-block and purified using MiniElute columns (Qiagen). Approximately 40 ng of insert DNA were ligated to 10 ng of *Bam*HI-digested pZERO-II vector with t4-ligase (Invitrogen). Into DHB10 ElectroMax *Escherichia coli* cells (Invitrogen), 0.5-µL aliquots were electroporated and they were then screened with kanamycin. The primary *B. stricta* library contained 12,000 colony-forming units (cfu) with an average insert size of approximately 5.0 kb and the primary *A. lyrata* library contained 4,000 cfu with an average insert size of approximately 5.5 kb.

End sequencing of genomic clones and comparison to *A. thaliana* genome

Nine hundred and sixty (960) genomic shotgun plasmid clones from both *A. lyrata* and *B. stricta* were end-

sequenced, resulting in two sequences per clone. Sequencing was performed using BigDye chemistry and bases were called with the *phred* algorithm in the *Phred/cross_match/Phrap* program (Ewing and Green 1998; Ewing et al. 1998). Vector sequences and low quality regions were then removed with the algorithm *cross_match* of the *Phred/cross_match/Phrap* program. A base-call was considered to be of high quality if it had a *phred* quality score of at least 30, and the five left and right neighboring base-calls had a *phred* score of at least 20. We also removed all sequences having similarity to plastid (i.e. chloroplast or mitochondrial) sequence.

In order to examine the influence of repetitive elements in our sequences, two bioinformatic filtering regimes were implemented. These filtering regimes differed principally in the inclusion versus exclusion in the resulting datasets of end-sequences having similarity to repetitive sequences. Thus, two datasets were derived from the full set of end-sequences, one for each filtering regime. Comparison of these two datasets allows us to infer the role of repetitive DNA in genome size changes (e.g. transposable elements, tandem repeats, etc.).

To compile the first dataset (hereafter “unfiltered”), BLASTn with default parameters was used to compare each set of end-sequences to itself in order to produce a non-redundant collection of inserts with paired end-sequences (Altschul et al. 1997). BLASTn (gap-opening penalty = 5, gap-extension penalty = 2, all other parameters set to default) was used to identify similarity between every non-redundant end-sequence and the *A. thaliana* nuclear genome (chromosome pseudomolecules NC_003070, NC_003071, NC_003074, NC_003075 and NC_003076; http://ftp.ncbi.nlm.nih.gov/genomes/Arabidopsis_thaliana). The resultant BLAST output was parsed, and end-sequences with at least one match to the *A. thaliana* nuclear genome with an *E* value greater than $10e^{-20}$ were scored as a “hit”; end-sequences that failed to meet this criterion were deemed as not having similarity to the *A. thaliana* nuclear genome (no hit).

To compile the second dataset (hereafter “filtered”), a number of additional bioinformatic filtering steps were applied. First, in order to avoid over-sampling regions of the genome resulting from the non-random cutting of the restriction enzyme, all sequence reads were clustered separately for each species with *CAP3* (Huang and Madan 1999) and the end-sequences from clones that formed clusters were discarded from further analysis. Second, the remaining end-sequences were compared against the *A. thaliana* genome with BLASTn (arabi all dna v170902) and scored as a “hit” if they had at least one match with an *E* value greater than $10e^{-20}$. End-sequences that did not meet this criterion and end-sequences for which the logarithm of the difference of the *E* value of the best and the

second best hit was less than 10 (to exclude repetitive sequences) were scored as “no hit”. The third and final step removed from further analysis end-sequences from clones for which the two end-sequences either matched to different chromosomes or spanned a region of more than 25,000 bp on the same chromosome of the *A. thaliana* genome.

Both bioinformatic filtering regimes resulted in three classes of clones: those from which both end-sequences passed all filtering steps (two-hit clones), those from which only one end-sequence passed all filtering steps (one-hit clones), and those from which neither end-sequence passed all filtering steps (zero-hit clones). For both datasets, we tested whether the two ends of the genomic clones were independent from one another using a chi-square test on the number of clones in each category. The chi-square test examined if the observed distribution of clones among these three categories was significantly different from the distribution expected under an assumption of independence between the two ends in their “hit” versus “no-hit” status.

Gel-based estimation of clone insert sizes

In order to conduct a conservative comparative analysis, only two-hit clones from the filtered dataset were used for pair-wise comparisons of clone insert sizes to their *A. thaliana* homolog. To measure the size of the insert, clones were digested to completion with *Mph1103I* (MBI Fermenta) and size-fractionated on a 0.8% agarose gel, together with size standards. Sizing of fragments was performed on an UV visualizer. To get the size of the homologous region in *A. thaliana*, the end-sequences of these clones were pair-wise aligned with their best matching *A. thaliana* BAC sequence using *cross_match*, which employs the Smith–Waterman algorithm for local pairwise alignments (Ewing et al. 1998; Smith and Waterman 1981). Using the alignment, the start and end positions of the query sequence and of the corresponding BAC clone were determined. These positions were then mapped onto the *A. thaliana* pseudochromosomes using the MIPS annotation (v170902).

Fully sequencing a s-set of 15 clones of *A. lyrata*

Inserts of 15 of the two-hit clones from the filtered dataset were selected for complete sequencing using the Gene-Jumper Primer Insertion Kit for Sequencing (Invitrogen). The two-hit clones were first assigned to one of three categories based on their size relative to their homologous *A. thaliana* region (larger, smaller, or the same size). Within each category, clones were randomly chosen. After electroporation into GeneHogs electrocompetent *E. coli* cells (Invitrogen), colonies were picked, grown, and the

plasmid isolated using standard procedures. Sequencing, base-calling, and low-quality sequence trimming were performed as described earlier. Sequences were assembled with *CAP3* and manually edited.

Sequences were compared with the *A. thaliana* genome using BLASTn. Where significant matches existed, sequences were pair-wise aligned with *A. thaliana* BACs using DOTTER (Sonnhammer and Durbin 1995) and VISTA (Bray et al. 2003; Couronne et al. 2003). Sequence annotation was performed manually by comparison of the sequences with homologous *A. thaliana* regions, as well as with the help of the GENSCAN gene prediction algorithm (Burset and Guigo 1996), *NetGene2* splice-site prediction on-line server (Brunak et al. 1991; Hebsgaard et al. 1996), and *RepBase* transposon database (Jurka et al. 2005).

Results

Phylogenetic analysis

The aligned matrix of all four markers was 3,376 bp in length, which comprised 491 (14.5%) parsimony informative characters. For the nuclear markers, the aligned matrix of *Chs* sequences was 706 bp long, of which 173 (24.5%) were parsimony informative, whereas the aligned matrix of ITS sequences was 651 bp in length, of which 165 (25.0%) were parsimony informative. Among our chloroplast markers, the aligned matrix of *matK* sequences was 1,517 bp long, of which 117 (7.7%) were parsimony informative, whereas the aligned matrix of *trnL* sequences was 502 bp long, of which 36 (7.2%) were parsimony informative. The genus *Aethionema* was used as an outgroup.

Parsimony analysis of the combined dataset recovered a single most-parsimonious tree with a length of 2,165 (CI = 0.624, RI = 0.533). In all four replicate Bayesian analyses of the combined dataset, the same topology was recovered with a log-likelihood score of approximately −16,260. The parsimony and Bayesian trees conflicted in the placement of two lineages, one leading to *Lepidium* and the other a clade consisting of the genera *Fourraea*, *Sisymbrium*, and *Eutrema*. The positions of these two lineages are reversed between the two analyses. The Bayesian topology, however, is only one parsimony step longer.

The topology recovered via Bayesian analyses was used in all subsequent analyses and is the tree shown in Fig. 1. Bayesian posterior probabilities are placed above branches leading to clades and parsimony bootstrap values below (Fig. 1). This tree, rather than the parsimony tree, was used because: (1) among the individual markers, only *Chs* supports a basal position for *Lepidium*; (2) topological tests could not reject the more nested placement of *Lepidium*

(indeed, there is only a one-step difference under parsimony); and, (3) another study using a different mix of markers supports a distant relationship between *Sisymbrium*, *Eutrema* (*Thellungiella*), and the “Halimolobine” Brassicaceae (Hall et al. 2002).

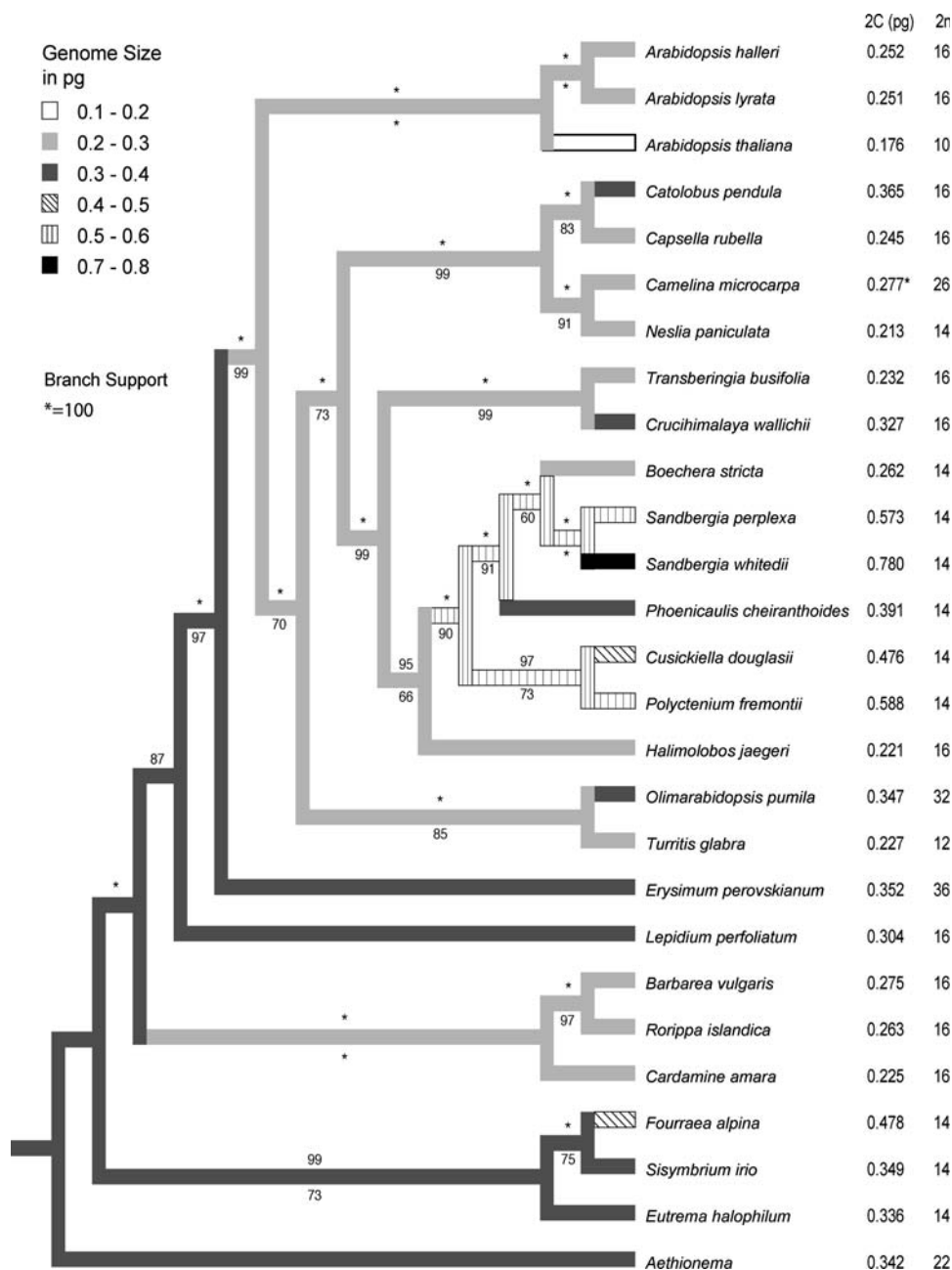
Genome size measurements

The genome size (2C) values for the taxa included in this study and their chromosome counts (2n) are listed in Table 1 and also shown in Fig. 1. Conversion of the continuous measurements of genome size (2C) into discrete

size categories resulted in nine possible character states (see Fig. 1). The small size of the *A. thaliana* genome is clearly revealed to be a derived state under a parsimony reconstruction of character evolution. Similarly, *B. stricta* also has a genome size that is smaller than its immediate relatives.

Reconstruction of the evolution of genome size (2C) in MacClade required 13 steps when the states were considered to be unordered and 17 when considered as ordered. These comprise six unambiguous transitions to a larger genome and seven unambiguous transitions to a smaller genome (see Fig. 1). Under our modified permutation test,

Fig. 1 Phylogeny recovered by Bayesian analysis. Values along branches are support values for the corresponding node with Bayesian posterior probabilities above and parsimony bootstrap values below. Genome sizes in pg (2C) and chromosome counts (2n) are given to the right of the taxa



this total number of transitions is not statistically different from what is expected if the character states were randomly distributed across the tree when the character is unordered. However, the 17 steps when the character was treated as ordered fall into the smallest 1.7% of a random distribution of possible steps in the modified permutation test, suggesting phylogenetic constraint. Analysis of the evolution chromosome numbers ($2n$) in MacClade revealed that the observed number of character state changes is lower than the average value under a random distribution, suggesting phylogenetic constraint. Under the permutation method in MacClade described earlier, this result was significant when the character was treated as unordered but not significant when the character was treated as ordered.

Our analyses of genome size ($2C$) and chromosome number ($2n$) as continuous characters in the program Continuous (Pagel 1997, 1999) were unable to reject the null-hypothesis that genome size and chromosome number each evolved under a constant-variance random walk model (Lamba not significantly different from 1; Model A not rejected). Furthermore, we failed to reject the null hypothesis that genome size ($2C$) and chromosome number ($2n$) evolved independently (i.e. not correlated; $\rho = 0.205$). However, whereas there was significant support for genome size having evolved under a model of punctuated equilibrium in which most change happened at the tips (Kappa = 0, Delta = 2.048), the evolution of chromosome number was found to be significantly dependent on branch length (Kappa = 3).

End sequencing of *Arabidopsis lyrata* and *Boechera stricta* genomic clones

The 960 clones we end-sequenced from each of the *A. lyrata* and *B. stricta* libraries resulted in a total of 3,840 sequence reads with a mean read length of 390 bp. The unfiltered dataset contained 619 clones from the *A. lyrata* library and 686 from the *B. stricta* library. The filtered dataset comprised 477 clones for *A. lyrata* and 568 for *B. stricta*. The BLASTn comparisons to *A. thaliana* (cutoff E value of 1^{-20}) revealed that a higher proportion of *A. lyrata* clones than *B. stricta* clones had a significant match (46 vs. 31%). The full complement of end-sequences (over 10 bp) was submitted to GenBank with accession numbers DX922572–DX924258 for *A. lyrata* and DX924259–DX926011 for *B. stricta*.

The additional quality trimming and filtering steps used to created the filtered dataset reduced the numbers of clones used for the analysis of segment lengths (i.e. two-hit clones) to 172 for *A. lyrata* and 150 for *B. stricta*. Almost all of the clones have the expected insert size range of 5–6 kb. Pairwise comparisons of the estimated sizes of the

inserts from these clones to the lengths of homologous regions in *A. thaliana* are shown in Fig. 2.

Both *A. lyrata* and *B. stricta* homologous regions are slightly but significantly larger compared with *A. thaliana* in a pairwise Wilcoxon Signed Rank test (Table 2). The mean and median for the size differences between *A. lyrata* clones and their homologous *A. thaliana* regions were, respectively, 396.9 and 205.5 bp larger (SD = 1,795.2). Meanwhile, the mean and median for the size differences between *B. stricta* clones and their homologous *A. thaliana* regions were, respectively, 82.7 and 277.5 bp larger (SD = 2,664.3). The distribution of comparative sizes in both species is unimodal but with a tail of smaller clone sizes (see Fig. 2). These measurements were not made for the set of two-hit clones in the unfiltered dataset because the ambiguity of homology to the highly similar regions in *A. thaliana* could make those values unreliable.

The distributions of the clones among the categories of two-hit (both end-sequences of a clone passed all filtering steps), one-hit (only one end-sequence passed all filtering steps), and zero-hit (neither end-sequence passed all filtering steps) were different between the two filtering regimes for both *A. lyrata* and *B. stricta* (Table 3). The numbers observed for the unfiltered dataset were significantly different from the expectations for all three categories under the chi-square test (*A. lyrata*: $\chi^2 = 124.13$, $P < 0.001$; *B. stricta*: $\chi^2 = 88.85$, $P < 0.001$) due to a deficit of one-hit clones. For the filtered dataset, the observed numbers were not significantly different from the expected according to the chi-square test (*A. lyrata*: $\chi^2 = 2.28$, $P = 0.320$; *B. stricta*: $\chi^2 = 0.06$, $P = 0.971$). Given that these two datasets differ mainly in the rigorousness with which they exclude repetitive sequences, comparing the results from these two bioinformatic regimes suggests that repetitive DNA is responsible for excesses in the zero-hit and two-hit categories for the unfiltered dataset. This pattern is predicted under a coarse indel model of genome size changes, where insertions and deletions >5 kb result in the hit-status of paired clone ends being correlated.

Fully sequenced clones

Complete sequencing of 15 of the two-hit clones from *A. lyrata* revealed that the average value by which the gel based estimation erred from the sequenced length was 59 bp (± 57 bp). In the clones sequenced, much of the size difference between the homologous *A. lyrata* and *A. thaliana* regions was found to be due to the presence of transposon-like and repetitive sequence in the larger homologous region (see supplementary data). The transposon-like sequences had similarity to a variety of Class I and Class II elements. For those sequences with similarity

Fig. 2 Graphs comparing sizes of *A. lyrata* and *B. stricta* inserts from two-hit clones in the filtered dataset to their homologous region in the *A. thaliana* genome. The top panels show the direct comparison of the sizes of the *A. lyrata* and *B. stricta* inserts from the genomic shotgun libraries to the sizes of their *A. thaliana* homologs. The bottom panels show the distribution of the size differences between the *A. lyrata* and *B. stricta* inserts to their *A. thaliana* homologs

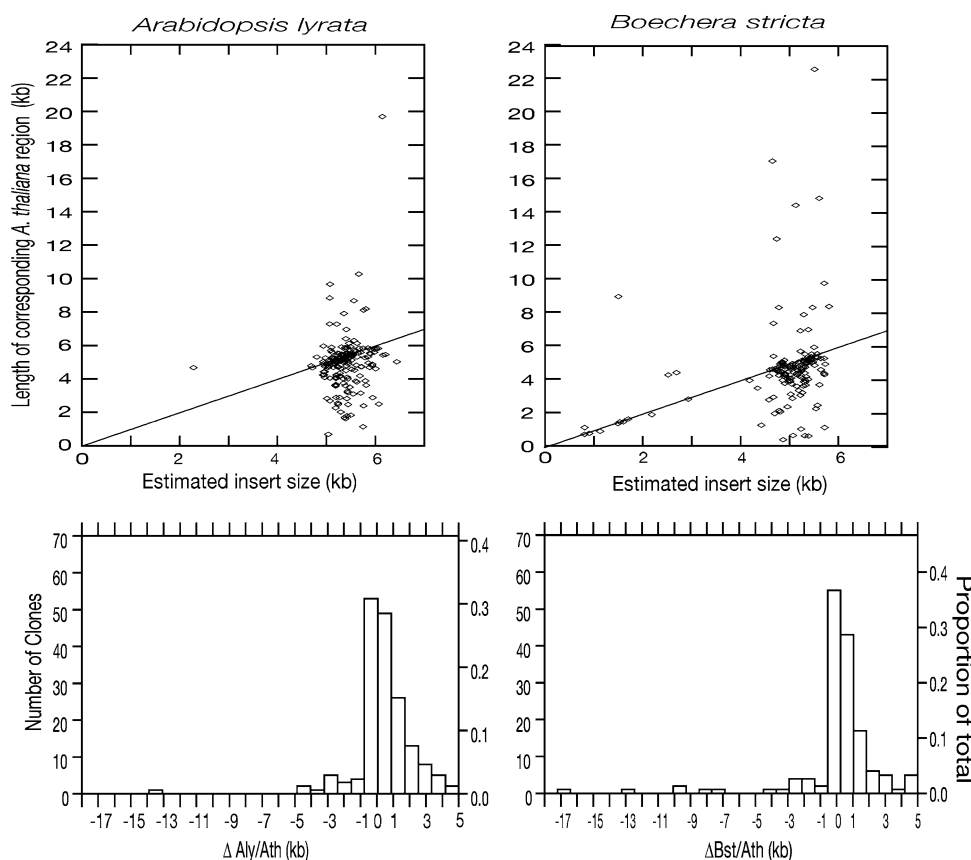


Table 2 Summary of the difference between insert sizes (reported as number of basepairs) and their homologous region in *Arabidopsis thaliana*

	<i>Arabidopsis lyrata</i>	<i>Boechera stricta</i>
Median ^a	205.5	277.5
Mean ^a	396.9	82.7
Minimum ^a	−13,594	−17,133
Maximum ^a	4,567	4,677
Standard deviation	1,795.2	2,664.3
<i>n</i>	172	150
Test statistics ^b		
<i>Z</i>	−4.874 ^c	−4.920 ^c
Sign. (two-tailed)	<0.0001	<0.0001

^a Values refer to the difference in length between insert from target species and homologous region in *A. thaliana* (i.e. Length of *A. lyrata* insert in bp—length of homologous region in *A. thaliana* in bp)

^b Wilcoxon Signed Rank test

^c Based on positive ranks

to retrotransposons (Class I), this included Ta1, subfamily Tf (LINE), other LINES, and non-LTRs. For those sequences with similarity to transposons (Class II), this included *En/Spm*, ARNOLDY, and MITE. Those clones that were similarly sized to their *A. thaliana* homolog exhibited a high sequence similarity between the two

species. These results are consistent with those observed by Windsor et al. (2006).

The length of exons and introns within genes of the sequenced *A. lyrata* clones were compared to their homologs in *A. thaliana*. This comparison included 13 complete genes and 22 incomplete genes for a total of 119 exons and 97 introns, excluding two introns (79 and 93 bp) found in *A. lyrata* but not in *A. thaliana* (Table 4). No significant difference in the length of either exons or introns was detected under a Wilcoxon Signed Ranks test. This suggests that most of the addition or deletion of genetic material occurred in intergenic regions. The complete annotated sequences of these clones were submitted to GenBank and have the accession numbers EU379001–EU379015.

Discussion

The use of four molecular markers (two nuclear and two chloroplast) and sampling among the genera thought closely related to *A. thaliana* provides a robust hypothesis of the phylogenetic relationships among these taxa. In addition, the mapping of genome sizes onto the phylogeny, and the subsequent examination of the mode, tempo, and directionality of genome size changes, provides a more critical assessment of the evolution of genome size than

Table 3 Numbers of clones of *Arabidopsis lyrata* and *Boechera stricta* having end-sequences that BLASTed to the *Arabidopsis thaliana* genome in the pattern indicated

	Zero hit clones	One hit clones	Two hit clones	Total number of clones	χ^2	<i>P</i> value
Unfiltered dataset						
<i>A. lyrata</i>	111 (89.2)	248 (291.6)	260 (238.2)	619	124.13	<0.001
<i>B. stricta</i>	224 (179.1)	277 (316.8)	185 (140.1)	686	88.85	<0.001
Filtered dataset						
<i>A. lyrata</i>	74 (75.3)	231 (228.4)	172 (173.3)	477	2.28	0.320
<i>B. stricta</i>	152 (143.0)	266 (284.0)	150 (141.0)	568	0.06	0.971

Values in parentheses are the expected values based on the observed frequency with which end-sequences “hit” the *A. thaliana* genome assuming that the hit/no-hit status of clone ends is not correlated

Categories correspond to those clones for which: neither end-sequence BLASTed to the *A. thaliana* genome (zero-hit); only one end-sequence BLASTed to the *A. thaliana* genome (one-hit); both end-sequences BLASTed to the *A. thaliana* genome (two hit)

Table 4 Comparison of exon and intron length within genes from the 15 fully sequenced two-hit clones from *A. lyrata* to their homologs in *A. thaliana*

	Mean length (bp)		Median length (bp)		Standard deviation		Sample size (<i>n</i>)	Wilcoxon Signed Rank test	
	<i>A. lyrata</i>	<i>A. thaliana</i>	<i>A. lyrata</i>	<i>A. thaliana</i>	<i>A. lyrata</i>	<i>A. thaliana</i>		Z test	<i>P</i> test
Exons	166	166 ^a	114	114	168.1	168.5	119	−0.090 ^c	0.928
Introns	153	151 ^b	93	93	180.4	170	97 ^c	−0.197 ^d	0.844

^a *Arabidopsis* genome average = ~250 bp (AGI, 2000)

^b *Arabidopsis* genome average = ~160 bp (AGI, 2000)

^c Based on negative ranks

^d Based on positive ranks

^e *A. lyrata* contained two introns (79, 93 bp) that were not present in *A. thaliana* and were therefore excluded from the analysis

heretofore available. Finally, comparisons of genomic clones from *A. lyrata* and *B. stricta* to *A. thaliana* provide an in-depth look at the underlying patterns of genome size evolution. We are thus able to not only describe the history of genome size evolution in this area of the Brassicaceae but also illustrate where patterns in our observed data are consistent with theoretical mechanisms generating diversity in genome size. This should provide a guide for further research on this topic.

Genome size changes across the phylogeny

The phylogeny recovered in this study (Fig. 1) is strongly supported and is similar to previous studies that have incorporated some of the taxa included here (Bailey et al. 2002; Hall et al. 2002; Johnston et al. 2005; Koch et al. 2001, 2003; O’Kane and Al-Shehbaz 2003). The one ambiguity is the placement of two basal lineages, which are swapped between the Bayesian and parsimony analyses. However, this does not affect our reconstruction of genome sizes since *Lepidium* and the lineage leading to *Fourraea*, *Sisymbrium*, and *Eutrema* have the same character state for genome size (Fig. 1).

The most parsimonious explanation for the small genome size of *A. thaliana* is a *reduction* in that lineage, not an *increase* in the sister clade (Fig. 1). This agrees with the conclusions of Johnston et al. (2005), although they found *Arabidopsis* to have a sister relationship with a clade containing *Capsella*, *Olimarabidopsis*, and *Crucihimalaya*. The differences in our topologies are likely due to our increased sampling of genera, denser taxonomic sampling near *Arabidopsis*, and the addition in our dataset of three other molecular markers. The genome size of 0.176 pg that we measured for *A. thaliana*, while slightly higher than the commonly reported value of 0.16 pg, is not much different from those reported by Johnston et al. (2005). Our analysis also indicated a genome size reduction in *B. stricta* relative to its sister taxa.

The results from our character mapping in MacCladeTM and Continuous find significant support for species-specific changes in genome size (2C). This is consistent with findings in other groups of organisms in which genome size changes occur along terminal branches (Grover et al. 2004; Neafsey and Palumbi 2003). In our MacCladeTM analyses, there is a greater tendency for genome size to increase (nine transitions) than for genome size to decrease (three

transitions) (Fig. 1), but this is likely due to the fact that there are more taxa with small (0.0–0.5 pg) genomes than with large (0.5–0.9 pg) genomes (22 vs. 5, respectively). Indeed, the failure to reject the null-hypothesis that genome sizes are distributed randomly across the tree under an unordered treatment of the character is consistent with no phylogenetic constraint. Although the result was significant when the character was treated as ordered, this was not as conservative a test. Our analyses using the program Continuous failed to support a model of directional evolution for either genome size (2C) or chromosome number (2n), but did find significant support for phylogenetic constraint in the evolution of both characters. The analyses in Continuous also found that the evolution of genome size (2C) and chromosome number (2n) is not correlated, and that these two characters appear to have different modes and tempos of evolution. The final picture suggests that genome size, at the scale we investigated, is free to increase and decrease over evolutionary time.

Mechanisms of change in genome size

An important distinction must be drawn in the evolution of genome size between the most common mechanistic event and the mechanism by which the majority of DNA addition or removal occurs. That is, a few events resulting in dramatic change of the total amount of DNA may overshadow the fact that most events have a very small effect on total genome size. In addition, mechanisms shown to work in the laboratory are not necessarily responsible for the variation in genome size that arises over the course of evolution. Nevertheless, our current understanding of genome size evolution does point toward a few generally important mechanisms. Furthermore, our data shed light on distinguishing between the most common mechanistic events and the events that likely account for most of the observed change in genome size among these lineages.

Two important mechanisms for genome size increases are polyploidization and the insertion of transposable elements (Bennetzen 2000; Kellogg and Bennetzen 2004). We do not address polyploidization in this paper but it is a relatively well understood phenomenon and evidence for multiple rounds of polyploidization in the evolutionary past of *A. thaliana* has already been documented (Simillion et al. 2002; Wendel 2000). Furthermore, polyploidy is known to occur at the tips of the Brassicaceae tree, for example in the genera *Arabidopsis* and *Boechera* (Clausen and Koch 2006; Schranz et al. 2005). Meanwhile, the importance of transposable elements in genome evolution is also well documented (Bartolome et al. 2002; Bennetzen 2000; Biemont and Vieira 2005; Kumar and Bennetzen 1999; Lenoir et al. 2005). Thus, it is not surprising that our sequenced clones revealed a correlation between the

presence of transposable elements and increases in the size of homologous regions (see online supplementary data associated with this article).

Mechanisms that might account for reductions in genome size across evolutionary time are not as obvious (Rabinowicz 2000). One proposed mechanism is the loss of whole chromosomes or genes following polyploidization events (Leitch and Bennett 2004). But our data reveal no correlation between chromosome number and genome size from an evolutionary perspective. Petrov (2002) and Petrov and Hartl (1997) have also suggested that biased mutation could lead to genome size reduction. And, evidence from Devos et al. (2002) indicates that “illegitimate recombination” is another means by which plants can avoid genomic obesity. In this scenario, breaks in double stranded DNA are repaired via non-homologous end joining, which leads to small-scale deletions (Devos et al. 2002; Gregory 2005a; Puchta 2005; Vitte and Panaud 2005). In the laboratory, DNA loss has been observed in *A. thaliana* during Double Strand Break repair (Kirik et al. 2000). Other conceivable mechanisms include unequal homologous recombination (Bennetzen et al. 2005), increases in the rate of deletions relative to insertions, or increases in the size of deletions relative to insertions.

Patterns from our genomic clones can help to identify which of the theoretical mechanisms were actually involved in genome size changes among the taxa studied here. For example, the majority of changes to genome size are small but there are also a considerable number of large increases and decreases (Fig. 2). This suggests that at least two processes are at work. Meanwhile, the similar characteristics of the *A. lyrata* and *B. stricta* datasets, suggests that the processes underlying the flux of genome sizes in these taxa are similar (Table 3). Although our data point toward certain explanations for the change in genome size as more probable than others, it is clear that the modest but significant size difference seen between our clones and the homologous regions from *A. thaliana* does not completely account for the change in overall genome size.

The contrasting results from the chi-square analyses of our genomic clone classes from the unfiltered and filtered datasets highlight the significant role of insertions and deletions larger than 5 kb containing repetitive sequence in genome size changes. Nevertheless, the results from the filtered dataset also suggest that a portion of the changes to genome size is made of up insertion or deletion events that are smaller than 5 kb. Thus, the analyses indicate that both coarse- (changes larger than 5 kb) and fine-grained models (changes smaller than 5 kb) are at work in the evolution of genome size. Consequently, our data paint a picture of genome size change in which much of the reduction or increase of genome size results from large

insertions/deletions involving repetitive sequences or transposable elements, but where most of the insertion/deletion events involve mechanisms by which relatively small amounts of DNA are inserted or deleted.

Further support for a fine-grained model comes from the many small insertion and deletion events observed in the fully sequenced two-hit clones of *A. lyrata*. This is congruent with the predictions of the “illegitimate recombination” model for small deletion sizes in LTR-regions (Bennetzen et al. 2005; Devos et al. 2002). In addition, the lack of significant size difference in exons or introns of the clones sequenced from *A. lyrata* relative to *A. thaliana* reveals that insertion and deletion events are generally intergenic (Table 4). This has also been observed in a previous study comparing *A. thaliana* and *A. lyrata* (Kusaba et al. 2001), *Drosophila* (Bartolome et al. 2002), and a study comparing the genomes of *A. thaliana* and *C. rubella* (Boivin et al. 2004).

Evidence in support of a coarse grained model, whereby relatively few changes have a large effect, can be found in the results from our chi-square analysis and our analysis using Continuous (Kappa = 0). This scenario agrees with suggestions that transposon activity may occur in bursts within specific lineages, thus leading to relatively abrupt increases in genome size (Vitte and Panaud 2005). This pattern is also consistent with general observations of dramatic genomic rearrangements and more rapid evolution following speciation events (Navarro and Barton 2003). However, the fact that our character mapping finds no support for genome size increases in the *A. lyrata* and *B. stricta* lineages and that the transposon-like sequences in our fully sequenced clones corresponded to a variety of transposon types, suggests that transposon infection and expansion, such as seen in maize (SanMiguel et al. 1998), is not responsible for the differences in genome size we observe.

Genome size reduction in the *A. thaliana* lineage

The possible causes, as opposed to the mechanisms, for a small genome size remain unclear (Charlesworth and Barton 2004; Petrov 2001). In the case of *A. thaliana*, genome size reduction is correlated with changes in other characteristics such as a shift to an annual lifestyle, unique within the genus. This lifestyle is characterized by more rapid development, which might have selected for a smaller genome size (Bennett et al. 1998; Charlesworth and Barton 2004). The shift to self-fertilization, also apomorphic along this lineage, would have led to a small effective population size that could have acted as a barrier to the establishment of transposable elements, thwarting a major mechanism of genome size increase (Lynch and Conery 2003). And, once a small genome size became established, it could itself have exerted a selective pressure for removal

of non-coding DNA, creating a positive feedback loop (Gregory 2005b).

The interplay of several of these mechanisms could account for the dramatic reduction in genome size in *A. thaliana*. The 3.1–9 million years age range for the lineage (Koch et al. 2000) encompasses the 6 million years half-life reported for LTR-retrotransposons in rice (Ma et al. 2004), meaning that transposable elements present at the time of divergence would have lost much of their ability to increase genome size over the time that has since elapsed. Meanwhile, the concomitant shift to a weedy, annual, self-fertilizing lifestyle in *A. thaliana* could have significantly impaired mechanisms that would otherwise have increased genome size, such as the spread of newly introduced transposons into the genome. Mechanisms that act to decrease genome size, however, would have remained unaffected. In this scenario, evolutionary pressure does not directly act to reduce genome size but merely hinders genome size increase.

The scenario of a reduced rate of transposon insertion but unhindered deletion has been found in other organisms. Neafsey and Palumbi (2003) find the probable explanation for the reduced genome size in the smooth pufferfish lineage to be a decline in the rate of large-scale insertions. The situation in *A. thaliana* may be, however, a little more complicated. Although a small effective population size could thwart the establishment of new transposable elements, it would also be expected to lower the efficacy of selection for the removal of transposable elements. However, this does not seem to have been the case in *A. thaliana*, relative to *A. lyrata*, which might be due to a population expansion following the evolution of selfing (Wright et al. 2002). In addition, *A. thaliana* seems to have a more complicated population structure than would be expected for a self-fertilizing annual, suggesting that its effective population size might not be as small as previously thought (Nordborg et al. 2005; Schmid et al. 2006).

Further genome–genome comparisons

We have a robust phylogeny that includes multiple species of *Arabidopsis* and closely related genera. This allows us to polarize the overall direction of genome size changes with confidence and claim that most of the processes leading to genome size reduction in *A. thaliana* occurred on the branch leading to that taxon. We also have genomic data from a close relative of *A. thaliana* (*A. lyrata*) and a more distant relative (*B. stricta*). This allows us to compare the micro-level changes in genome composition among the three taxa and claim that multiple mechanisms are driving the evolution of genome size. There are, however, two caveats. The first is that since only the genome of

A. thaliana has been fully sequenced, we do not have a true three-way comparison between *A. thaliana*, *A. lyrata* and *B. stricta* and thus cannot polarize the individual indel events identified in our fully sequenced genomic clones. The second is that we do not have sufficient population or species level sampling to consider inter- versus intraspecific variation in genome size. Thus, confirmation of our claims will require greater population level measurements of genome size for the numerous Brassicaceae taxa in our phylogeny and direct comparisons of the complete genome sequences of the three taxa.

The sequencing of the *A. lyrata* genome will help to address the problem of polarizing the indel events by making direct comparisons between *B. stricta* and *A. lyrata* possible. It would also provide the potential to identify the transposons responsible for genome size changes. However, as was discovered with *Drosophila*, the whole-genome shotgun sequencing strategy may not be able to accurately place repetitive elements or indels, and thus it will be difficult to use the data generated to infer processes of genome size evolution (Benos et al. 2001). This means that even after the completion of this project, and the effort to sequence the genome of *Capsella*, the role of repetitive sequences in the evolution of the *A. thaliana* genome may remain unclear.

Conclusion

The summary picture that emerges from this study is that the small size of the *A. thaliana* genome is derived relative to its sister taxa, and that this is likely due to both large, coarse-grained insertion/deletion events and smaller, fine-grained insertion/deletion events. Furthermore, our data implicate transposable elements in the insertion events leading to increases in genome size or at least the maintenance of a larger genome size. We also found evidence that insertion/deletion events occurring in a fine-grained way happen in intergenic regions scattered across the genome. This study illustrates how we can extract much more from the data generated by genomics techniques when we integrate them into a comparative study that includes closely related non-model species.

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